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PATENT SPECIFICATION

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(54) IMPROVED PROCESS FOR CATALYTICALLY DEWAXING GAS OIL

(71) We, MOBIL OIL CORPORATION, a Corporation organised under the laws of the State of New York, United States of America, of 150 East 42nd Street, New York, New York, 10017, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be

particularly described in and by the following statement:-

This invention relates to a process for dewaxing petroleum oils and fractions thereof by selectively removing normal paraffinic and other undesirable hydrocarbons from petroleum oils in which they are present in admixture with other hydrocarbons, in order to lower the pour point of such oils. More particularly, the invention relates to an improved single-step process for selectivity removing normal paraffinic and other undesirable hydrocarbons from petroleum oils by contacting same with a specific zeolite catalyst in the presence of a large amount of co-fed water.

It is well known in the art to form various lubricating oils from hydrocarbon fractions derived from petroleum crudes. A heretofore practiced common procedure known in the art is to extract these hydrocarbon fractions with various solvents so as to give a raffinate of a desired high viscosity index, such material being resistant to changes in viscosity with changes in temperature and thus being useful under varying operating conditions. Moreover, it is particularly desired that the lube oil have a low pour point so that it can be effectively used at low temperature conditions, since excessive thickening at low temperature is often unacceptable. It is also known in the art to carry out dewaxing operations by contacting hydrocarbon fractions with crystalline aluminosilicate zeolites having pore sizes of about 5 Angstrom units so as to selectively remove normal paraffins.

The present invention is concerned with an improved process for dewaxing normal paraffin-containing oils which is more economical than conventional solvent dewaxing procedures or catalytic dewaxing procedures involving 5 Angstrom unit zeolites and which, with certain feedstocks, produces a higher product yield with equivalent or higher pour point

reduction and prolonged catalyst cycle time between regenerations.

Trace amounts of water used to promote various catalytic reactions, not including gas oil dewaxing, is known in the art. For example, U.S. Patent 3,546,100 teaches restricting the partial pressure of water in contact with a hydrocracking catalyst during hydrocracking of a hydrocarbon compound to within the range of 10 to 130 mm. U.S. Patent 3,649,524 teaches a

high temperature reforming process using only 8-20 ppm water.

Somewhat larger amounts of water have been used in high temperature catalytic dehydrogenation of hydrocarbons (U.S. Patent 3,907,921), wherein 25-3000 ppm of water is used; and in hydrodesulfurization of gas oils and cycle oils (U.S. Patent 3,720,602) wherein 5 to 50 percent of water is injected. In the latter patent, the catalyst material used does not include zeolite materials.

The use of large amounts of water, i.e. about 0.1 to about 15 moles water/mole hydrocarbon feedstock, would be expected, based upon the teaching of the art, to destroy

conventional porous, siliceous heterogeneous catalysts, such as used in the present method.

U.S. Patent 3,755,138 discloses a two-step process for dewaxing hydrocarbon oil feedstocks boiling above about 650°F having an intermediate pour point. One step of the patent process comprises solvent dewaxing and the other step comprises contact with a ZSM-5 type of zeolite in the absence of co-fed water. U.S. Patent Re. 28,398 discloses dewaxing a hydrocarbon oil boiling above 350°F by shape selective cracking and

	hydrocracking over a zeolite of ZSM-5 type without co-fed water. This invention relates to improved processing of gas oils wherein said processing comprises contacting a gas oil feedstock in the presence of from about 0.1 to about 15 moles of co-fed	
5	water/mole of gas oil feedstock with a catalyst comprising a crystalline aluminosilicate zeolite characterized by a silica/alumina mole ratio of at least 12 and a constraint index, hereinafter defined, within the approximate range of 1 to 12.	5
	The feedstock intended for treatment in accordance with the present invention may be generally defined as hydrocarbon oils boiling above about 350°F and particularly between about 350°F and about 1,100°F. Processing of such feedstocks having a high nitrogen content	
10	of from about 0.01 to about 3 wt. percent is most advantageously benefited by the present improved processing technique.	10
15	The crystalline aluminosilicate zeolites used in the catalyst composition of the process of this invention are referred to generally as ZSM-5 type or as behaving like ZSM-5 and include ZSM-5, ZSM-11, ZSM-12, ZSM-35 and ZSM-38, more particularly described hereinafter. The catalyst composition useful in this invention comprises a crystalline aluminosilicate zeolite characterized by a silica/alumina mole ratio of at least 12 and a constraint index of	15
	from about 1 to about 12, non-limiting examples of which include ZSM-5, ZSM-11, ZSM-12, ZSM-35 and ZSM-38. Zeolite ZSM-5 is taught by U.S. Patent 3,702,886. In a preferred synthesized form, the	
20	zeolite ZSM-5 for use in the catalyst composition useful in this invention has a formula, in terms of mole ratios of oxides in anhydrous state, as follows: $(0.9 \pm 0.2) \text{ M}_2\text{O}: \text{Al}_2\text{O}_3: \text{xSiO}_2$	20
25.	wherein M is selected from a mixture of alkali metal cations, especially sodium, and tetraalkylammonium cations, the alkyl groups of which preferably contain 2 to 5 carbon atoms, and x is at least 5. Particularly preferred is a zeolite having the formula in the anydrous state as follows:	25
	$(0.9 \pm 0.2) \frac{M_2O}{n} : Al_2O_3 : ZSiO_2$	
30	wherein Z is from greater than 30 to about 350 or higher. Zeolite ZSM-11 is taught by U.S. Patent 3,709,979. In the as synthesized form, the zeolite ZSM-11 for use in the catalyst composition useful in this invention has a formula, in terms of mole ratios of oxides in the anhydrous state, as follows:	30
35	$(0.9 \pm 0.3) \frac{M_2O}{n}$: 20 to 90 SiO ₂	35
	wherein M is a mixture of at least one of the quaternary cations of a Group V-A element of the Periodic Table and alkali metal cations, especially sodium. The original cations can be present to that the amount of quaternary metal cations is between 10 and 90 percent of the total amount of the original cations. Thus, the zeolite can be expressed by the following	· ·.
40	formula in terms of mole ratios of oxides: $(0.9 \pm 0.3) (xXR_4 + 1-xM_2O) : Al_2O_3 : 20 \text{ to } 90 \text{ SiO}_2$	40
45	wherein R is an alkyl or aryl group having between 1 and 7 carbon atoms, M is an alkali metal cation, X is a group V-A element, especially a metal, and x is between 0.1 and 0.9. Zeolite ZSM-12 is taught by U.S. Patent 3,832,449.	. 45
	ZSM-35 is described in U.S. Patent No. 4,016,245. This zeolite can be identified, in terms of mole ratios of oxides and in the anhydrous state, as follows: (0.3 - 2.5)R ₂ O: (0 - 0.8)M ₂ O: Al ₂ O ₃ : xSiO ₂	• :
50	wherein R is an organic nitrogen-containing cation derived from ethylenediamine or	50
	In a preferred synthesized form, zeolite ZSM-35 has a formula, in terms of mole ratios of oxides and in the anhydrous state, as follows: (0.4 - 2.5)R ₂ O: (0 - 0.6)M ₂ O: Al ₂ O ₃ : ySiO ₂	
55	wherein R is an organic nitrogen-containing cation derived from ethylenediamine or pyrrolidine, M is an alkali metal, especially sodium, and y is from greater than 8 to about 50. ZSM-38 is described in U.S. Patent No. 4,046,859. This zeolite can be identified, in terms	55
60	of mole ratios of oxides and in the anhydrous state, as follows: (0.3 - 2.5)R ₂ O: (0 - 0.8)M ₂ O: Al ₂ O ₃ : xSiO ₂ wherein R is an organic nitrogen-containing cation derived from a 2-(hydroxyalkyl)	60
	trialkylammonium compound, x is greater than 8 and M is an alkali metal cation, and is characterized by a specified X-ray powder diffraction pattern. In a preferred synthesized form, the zeolite has a formula, in terms of mole ratios of oxides	
65	and in the anhydrous state, as follows: (0.4 - 2.5)R ₂ O: (0 - 0.6)M ₂ O: Al ₂ O ₃ : ySiO ₂	65

wherein R is an organic nitrogen-containing cation derived from a 2-(hydroxyalkyl) trialkylammonium compound, wherein alkyl is methyl, ethyl or a combination thereof, M is an alkali metal, especially sodium, and y is from greater than 8 to about 50. Although the zeolites herein described have unusually low alumina contents, i.e. high silica to alumina ratios, they are very active even when the silica to alumina ratio exceeds 30. The activity is surprising since catalytic activity is generally attributed to framework aluminium atoms and cations associated with these aluminum atoms. These catalysts retain their crystallinity for long periods in spite of the presence of steam at high temperature which induces irreversible collapse of the framework of other zeolites, e.g. of the X and A type.

10 Furthermore, carbonaceous deposits, when formed, may be removed by burning at higher than usual temperatures to restore activity. In many environments the zeolites of this class exhibit very low coke forming capability, conducive to very long times on stream between burning regenerations. An important characteristic of the crystal structure of the zeolites for use herein is that they provide constrained access to, and egress from, the intracrystalline free space by virtue of having a pore dimension greater than about 5 Angstroms and pore windows of about a size such as would be provided by 10-membered rings of oxygen atoms. It is to be understood, of course, that these rings are those formed by the regular disposition of the tetrahedra making up the anionic framework of the crystalline aluminosilicate, the oxygen atoms themselves being bonded to the silicon or aluminum atoms at the centers of the tetrahedra. Briefly, the preferred type catalysts useful in this invention possess, in combination: a silica to alumina ratio of at least about 12; and a structure providing constrained access to the crystalline free space. The silica to alumina ratio referred to may be determined by conventional analysis. This ratio is meant to represent, as closely as possible, the ratio in the rigid anionic framework of 25 the zeolite crystal and to exclude aluminum in the binder or in cationic or other form within the channels. Although catalysts with the silica to alumina ratio of at least 12 are useful, it is preferred to use catalysts having higher ratios of at least about 30. Such catalysts, after activation, acquire an intracrystalline sorption capacity for normal hexane which is greater than that for water, i.e. they exhibit "hydrophobic" properties. It is believed that this hydrophobic character is advantageous in the present invention. The type zeolites useful in this invention freely sorb normal hexane and have a pore dimension greater than about 5 Angstroms, or, if elliptical in pore shape, at least the size of the pores in ZSM-5. In addition, the structure must provide constrained access to larger molecules. It is sometimes possible to judge from a known crystal structure whether such constrained access exists. For example, if the only pore windows in a crystal are formed by 8-membered rings of oxygen atoms, then access to molecules of larger cross-section then normal hexane is excluded and the zeolite is not of the desired type. Windows of 10-membered rings are preferred, although, in some instances, excessive puckering or pore blockage may render these catalysts ineffective. Twelve-membered rings do not generally 40 appear to offer sufficient constraint to produce the advantageous conversions. Also, structures can be conceived due to pore blockage or other cause, that may be operative. Rather than attempt to judge from crystal structure whether or not a catalyst possesses the necessary constrained access, a simple determination of the "constraint index" may be made by passing cotinuously a mixture of an equal weight of normal hexane and 3-methylpentane over a small sample, approximately 1 gram or less, of catalyst at atmospheric pressure according to the following procedure. A sample of the catalyst, in the form of pellets or extrudate, is crushed to a particle size about that of coarse sand and mounted in a glass tube. Prior to testing, the catalyst is treated with a strem of air at 1000°F for at least 15 minutes. The catalyst is then flushed with helium and the temperature adjusted between 550°F to give an overall conversion between 10% and 60%. The mixture of hydrocarbons is passed at I liquid hourly space velocity (i.e., 1 volume of liquid hydrocarbon per volume of catalyst per hour) over the catalyst with a helium dilution to give a helium to total hydrocarbon mole ratio of 4:1. After 20 minutes on stream, a sample of the effluent is taken and analyzed, most conveniently by gas chromatography, to determin the fraction remaining unchanged for each of the two hydrocarbons.

log 10 (fraction of 3-methylpentane remaining) The constraint index approimates the ratio of the cracking rate constants for the two hydrocarbons. Catalysts suitable for the present invention are those having a constraint index in the approximate range of 1 to 12. Constraint Index (CI) values for some typical catalysts, including those useful herein, are:

The "constraint index" is calculated as follows: Constraint Index = log10 (fraction of n-hexane remaining)

determined by classical pyknometer techniques. For example, it may be determined by

immersing the dry hydrogen form of the zeolite in an organic solvent which is not sorbed by the crystal. It is possible that the unusual sustained activity and stability of this class of zeolite is associated with its high crystal anionic framework density of not less than about 1.6 grams per cubic centimeter. This high density of course must be associated with a relatively small amount of free space within the crystal, which might be expected to result in more stable structures. This free space, however, is important as the locus of catalytic activity.

Crystal framework densities of some typical zeolites are:

10:	•	•	•		10
10		Zeolite	Void Volume	Framework Density	10
		Ferrierite	0.28 cc/cc	1.76 g/cc	1.5
15		Mordenite	.28	1.7	15
		ZSM-5, -11	.29	1.79	•
20		Dachiardite	.32	1.72	20
		L	.32	1.61	
		Clinoptilolite	.34	1.71	. 25
25	٠.	Laumontite	.34	1.77	25
		ZSM-4	.38	1.65	
30		Heulandite	.39	1.69	30
· :		P	.41	1.57	· ·
	•	Offretite	.40	1.55	. 25
35		Levynite	.40	1.54	35
•		Erionite	.35	1.51	
40	•	Gmelinite	.44	1.46	. 40
•	· .	Chabazite .	.47	1:45	•
		A	.5	1.3	45
45 .		Y	.48	1.27	

Members of the above group of zeolites for use in the catalyst composition of the present invention possess definite distinguishing crystalline structures as evidenced by the above U.S. Patents incorporated herein by reference.

The synthetic ZSM-35 zeolite possessess a definite distinguishing crystalline structure whose X-ray diffraction pattern shows substantially the significant lines set forth in Table 1.

TABLE 1

•	Interplanar Spacing	Relative Intensity	
5	9.6 ± 0.20	Very Strong-Very, Very Strong	5
	7.10 ± 0.15	Medium	
10	6.98 ± 0.14	Medium	
10	6.64 ± 0.14	Medium	10
	5.78 ± 0.12	Weak	
	5.68 ± 0.12	Weak	. :
15	4.97 ± 0.10	Weak	15
	4.58 ± 0.09	Weak	
	3.99 ± 0.08	Strong	•
. 20	3.94 ± 0.08	Medium-Strong	20.
20	3.85 ± 0.08	Medium	20.
	3.78 ± 0.08	Strong	•
	3.74 ± 0.08	Weak	
25	3.66 ± 0.07	Medium	25
	3.54 ± 0.07	Very Strong	
	3.48 ± 0.07	Very Strong	
30	3.39 ± 0.07	Weak	30
	3.32 ± 0.07	Weak-Medium	•
	3.14 ± 0.06	Weak-Medium	•
35	2.90 ± 0.06	Weak	35
33	2.85 ± 0.06	Weak	
- +	2.71 ± 0.05	Weak	•
40	2.65 ± 0.05	Weak	40
40	2.62 ± 0.05	Weak	40
	2.58 ± 0.05	Weak .	
•	2.54 ± 0.05	Weak	٠
45	2.48 ± 0.05	Weak	45

The synthetic ZSM-38 zeolite possesses a definite distinguishing crystalline structure whose X-ray diffraction patterns shows substantially the significant lines set forth in Table 50 1A.

TABLE 1A

•	Interplanar Spacing	***	Relative Intensity	
. 5	9.8 ± 0.20		Strong	5
	9.1 ± 0.19		Medium	
٠.	8.0 ± 0.16		Weak	
10	7.1 ± 0.14		Medium	10
	6.7 ± 0.14		Medium	
	6.0 ± 0.12	, s	Weak	
15 ⁻	4.37 ± 0.09		Weak	16
: 13	4.23 ± 0.09		Weak	15
	4.01 ± 0.08		Very Strong	
••	3.81 ± 0.08	$(x,y) = \sup_{x \in \mathcal{X}} \{x,y \in \mathcal{Y} \mid x \in \mathcal{X}\}$	Very Strong	
20	3.69 ± 0.07		Medium	20
	3.57 ± 0.07		Very Strong	
	3.51 ± 0.07		Very Strong	
25	3.34 ± 0.07		Medium	25
•	3.17 ± 0.06		Strong	
	3.08 ± 0.06	• • •	Medium	
30	3.00 ± 0.06		Weak	30
	2.92 ± 0.06	•	Medium	
	2.73 ± 0.06		Weak	
 25'	2.66 ± 0.05	• • •	Weak	25
35	2.60 ± 0.05		Weak .	35 °
	2.49 ± 0.05		Weak	
		•		
40	These values were determined by s doublet of copper, and a scintillation coused. The peak heights, I, and the posit Bragg angle, were read from the spectro	inter spectrometer vions as a function o	with a strip chart pen recorder was f 2 times theta, where theta is the	40
45	I/I ₀ , where I ₀ is the intensity of the stron in Angstrom units, corresponding to understood that these X-ray diffraction above respectively identified zeolites. I substantially the same pattern with som	gest line or peak, an the recorded lines n patterns are chara lon exchange of the	idk (obs.), the interplanar spacing s, were calculated. It should be acteristic of all the species of the e sodium ion with cations reveals	. 45 .
50	relative intensity. Other minor variatic ratio of the particular sample, as well Zeolites ZSM-5, ZSM-11 and ZSM-1 as indicated in their respective patents	ons can occur deper as if it has been su 2 for use in the proo	nding on the silicon to aluminum bjected to thermal treatment. cess of this invention are prepared	50
55	Zeolite ZSM-35 can be suitably prep alkali metal oxide, preferably sodium o of aluminum, an oxide of silicon and wa of oxides, falling within the following	ared by preparing a xide, an organic nit iter and having a co	solution containing sources of an trogen-containing oxide, an oxide	55

	•	\cdot			
		TABLE 2			
		Broad	Preferred	•	
5	R ⁺	0.02 1.0			5
	$R^+ + M^+$	0.02 — 1.0	0.3 – 0.9		
10	OH7SiO ₂	0.05 - 0.5	0.07 - 0.49		10
	H ₂ O/OH ⁻	41 - 500	100 - 250		10
-	SiO ₂ /Al ₂ O ₃	8.8 - 200	12 - 60		
15			• • • • • • • • • • • • • • • • • • • •		15
20 25	wherein R is an organic nitrogethylenediamine and M is an alkali nizeolite are formed. (The quantity calkali without any organic base contiliquid and recovered. Typical react mixture to a temperature of from a about 6 hours to about 100 days. An about 400°F with the amount of time to about 80 days. The digestion of the gel particles separated from the reaction medium and water washing. The crystalline hours. Zeolite ZSM-38 can be suitably p	netal ion, and main of OH ⁻ is calculate ribution). Thereaftion conditions con bout 90°F to about 90°F to about eat a temperature is is carried out until as by cooling the product is dried, of	taining the mixture uned only from the inorter, the crystals are selsist of heating the fout 400°F for a period perature range is from such range being from the crystals form. The whole to room tempers, at 230°F, for from	atil crystals of the ganic sources of parated from the regoing reaction of time of from about 150°F to mabout 6 hours solid product is erature, filtering mabout 8 to 24	20
	alkali metal oxide, preferably sodius of aluminum, an oxide of silicon and of oxides, falling within the following	m oxide, an organi I water and having	c nitrogen-containing	oxide, an oxide	
35	•	TABLE 3			35
		Broad	Preferred		
40	. R+				40
	$R^+ + M^+$	0.2 - 1.0:	0.3 - 0.9		•
	OH7SiO ₂	0.05 - 0.5	0.07 - 0.49		
45	H ₂ O/OH ⁻	41 - 500	100 - 250	•	45
	SiO ₂ /Al ₂ O ₃	8.8 - 200	12 - 60	. •	•
50		•	•		50
55	wherein R is an organic nitroger trialkylammonium compound and M crystals of the zeolite are formed. (T sources of alkali without any org separated from the liquid and recov foregoing reaction mixture to a temp of time of from about 6 hours to about 150°F to about 400°F with the from about 6 hours to about 80 day. The digestion of the gel particles	I is an alkali metal interpretable definition of OH' anic base contributed. Typical reactorist 100 days. A more amount of time bys.	ion, and maintaining to scalculated only froution). Thereafter, to tion conditions consistent 90°F to about 40 perferred temperatuat a temperature in s	the mixture until om the inorganic the crystals are st of heating the 0°F for a period ure range is from uch range being	55
65	separated from the reaction medium and water washing. The crystalline p to 24 hours. For the improved process of this	, as by cooling the roduct is thereafte	whole to room temper dried, e.g. at 230°F1	erature, filtering for from about 8	65

combination with a support or binder material such as, for example, a porous inorganic oxide support or a clay binder. Non-limiting examples of such binder materials include alumina, zirconia, silica, magnesia, thoria, titania, boria and combinations thereof, generally in the form of dried inorganic oxide gels and gelatinous precipitates. Suitable clay materials include, 5 by way of example bentonite and kieselguhr. The relative proportion of suitable crystalling aluminosilicate zeolite of the total composition of catalyst and binder or support may vary widely with the zeolite content ranging from between about 30 to about 90 percent by weight and more usually in the range of about 50 to 80 percent by weight of the composition. Operating conditions employed in the process of the present invention are critical. Such conditions as temperature, pressure, space velocity, molar ratio of co-fed water to gas oil 10 feedstock, absence or presence of added hydrogen, and the presence of any diluents will have important effects on the process. The process of this invention is conducted in the liquid or mixed gas-liquid phase and with or without added hydrogen by contact in a reaction zone, such as, for example, a fixed bed of catalyst composition, under conversion effective conditions, said catalyst composition, under conversion effective conditions, said catalyst composition being characterised, as synthesized, as comprising one or more of the above-defined zeolite compositions. This process may be conducted in either batch or fluid bed operation with attendent benefits of either operation readily obtainable. The present improved process may be carried out at a temperature of between about 450°F and about 800°F, preferably from about 500°F to about 750°F, and at pressures ranging from about 50 psig up about 3000 psig, preferably from about 100 psig to about 1000 psig. The liquid hourly space velocity (LHSV) may be maintained at from about 0.1 hr⁻¹ to about 2 hr-1. Hydrogen circulation may be maintained at from 0 to about 10,000 scf/bbl. The preferred amount of co-fed water is from about 0.1 to about 5 moles of water/mole of gas oil feedstock. Within these limits the conditions of temperature and pressure will vary considerably depending upon equilibrium considerations and exact feed material. Optimum conditions are those in which maximum yields of desired dewaxed products are obtained and hence temperature and pressure will vary within a range of conversion levels designed to provide the highest selectivity and maximum yield. The starting feed materials for the present improved process are petroleum stocks boiling above about 350°F and containing straight chain and slightly branched chain hydrocarbons which selectively converted utilizing a catalyst composition as hereinabove particularly described. The improvement realized by way of the present process is substantial in that coking and aging rates of the catalyst are controlled while conversions remain favorable. The zeolite catalysts above defined for use herein are found to be hydrophobic and unique in their ability to uitlize the large molar equivalents of co-fed water to resist coking and aging at the activity levels maintained in the present process. The following specific examples will serve to illustrate the process of the present invention, without unduly limiting same. Examples 1 and 8 illustrate the preparation of suitable catalysts. Examples 2. 4 and 7 are incuded for comparison. Example 1 A silicate solution containing 90.9 pounds Q-Brand sodium silicate (8.8 wt. % Na₂O, 28.5 wt. % SiO2 and 62.7 wt. % H2O). 52.6 pounds H2O, and 118 grams Daxad 27 dispersant (sodium salt of polymerized substituted benzoid alkylsulfonic acid combined with an inert inorganic suspending agent) was mixed in a mixing nozzle with an acid solution containing 1430 grams Al₂(SO₄)₃, 14 H₂O, 3440 grams H₂SO₄, 4890 grams NaCl, and 54 pounds H₂O to form a gel which was discharged into a 30 gallon autoclave to which 1180 grams of H2O has 50 been previously added. The gel was whipped by agitation and 2840 grams of NaCl was added and thoroughly blended. The agitation was stopped and an organic solution containing 2780 grams tri-n-propylamine. 2390 grams n-propyl bromide, and methyl ethyl ketone was added as a layer on top of the gel. The autoclave was sealed and heated to about 220°F without agitation and held there for 14-15 hours to prereact the organics. At the end of the prereaction period the agitation was commenced at 90 rpm to start the initial crystallization period. After about 75-80 hours, the temperature was raised to 320°F and held there for about 3 hours to complete crystallization. The excess or unreacted organics were flashed off and the contents of the autoclave were cooled and discharged. Chemical analysis of the washed crystalline product was 2.21 wt.% Al₂O₃. 94.9 wt. % SiO₂. 0.81 wt. % Na. 0.67 wt. % N. and 8.2 wt. % carbon, and it was identified by X-ray as ZSM-5. After drying at about 250°F, the zeolite was mixed with alpha-alumina monohydrate and

 H_2O (65% zeolite. 35% alumina binder on ignited basis), then extruded to form 1/16 inch pellets. The pellets were calcined in nitrogen atmosphere for 3 hours at 1000°F, ion exchanged with 1 normal NH₄Cl at room temperature for 1 hour using 5 milliliters solution

35

per gram of dry zeolite, water-washed, and finally calcined in 100% air for 3 hours at 1000°F. Examples 2-7

The catalyst material prepared in Example 1, sized to 30-60 mesh, was charged to a 5/16th inch I.D. stainless steel tubing reactor with provision for water injection at the reactor inlet 5 from a positive displacement pump. Six separate runs were conducted in this reactor, each with 5 or 10 cc of fresh catalyst, under varying conditions and for different times on stream. The feedstock to the reactor was a 650-760°F Arab Gas Oil having a pour point of 55°F and a specific gravity of 0.8866. No hydrogen was added to the reactor for these runs. Reaction conditions and results are reported hereinafter in Table 4.

From the information generated by Examples 2-7, it is observed that aging rates were high at the reaction pressures used when water was not co-fed to the reactor, The beneficial affect of the large amounts of co-fed water on coke lay-down is clear by comparison of the examples run with and without water injection.

					•	
Example	7	M	4	٠.		:
December Dein	. 05		200	200	200	200
riessule, rag	1.0	1.0	1.0	1.0	٠:	1.1
Gas out, Lates	, ,	90.0	t ·	90.0		0.83
ngo, Linav	·. . 1	2.6	· 1	2.6		32.5
Mole ratio of n2 O/gas off	610	230	540	550		230
Initial temperature, F	820	830	770	. 09/		610
rinai temperature, 1. Timo on titonm dave	S .	7	∞	22	,	14-
Inition on stream, days.	, 1	34	29	. 10		9
Aguig laic, 1/uay	22.0	15.7	11.5	17.3	9.6	7.5

For 330°F product, pour point of 0°F.

Example 8

	•	Exa	тріе в			
5	A portion of the dried am with 1 normal Ni(NO ₃) ₂ at 1! extrudate, water-washed, dri final product was analyzed an	90°F for 4 hours ed, and finally c d found to conta	using 5 cc of exchange alcined in 100% air fo	solution per gram of dry	٠.	
10	A 3.3 cc portion of fresh charged to the reactor used in Examples 2-7. The catalyst was added to the reactor for	for Examples 2- as sulfided <i>in si</i> this run at 2500	7 for a run with the samue with a H ₂ S/H ₂ mixt of scf/bbl pressure was	me feedstock as used for ure at 750°C. Hydrogen	. 10	
15	LHSV of the gas oil feedstoc feedstock was 6.5 ± 1.0. Th Regeneration #1 Hydrogen regeneration a scf/bbl.	e catalyst was r	egenerated twice duri	ng the run as follows:	15	
20	Regeneration #2 Hydrogen regeneration at 500 psig and 625°F for 2 hours and 900°F for 18 hours at a H ₂ flow of 2500 scf/bbl. Results of the run proved that the use of co-fed water and hydrogen worked synergistically to prevent the catalyst from deactivating by coke formation and/or nitrogen compound deposition. Aging data for this example is summarized in Table 5.					
25		TA	BLE 5		- 25	
30	Cycle	Fresh	After Regen. #1	After Regen. #2		
	Days on stream	24	20	4	30	
35	Total days on stream	24	44	48	35	
	Initial temp., °F*	540	580	~ 580		
40	Final temp., °F*	660	640	-	40	
45	Aging rate, °F/day	5	3	-	45	
	*For 330°F ⁺ product, pour point of 0°F.					
50	WHAT WE CLAIM IS:-				50	
55	1. A catalytic process for feedstock with a catalyst silica/alumina mole ratio of at 1 to 12, in the presence of feedstock.	comprising a c	crystalline aluminosili onstraint index within t	cate zeolite having a	55	
60	 The process of Claim 1 The process of any on temperature from about 450° 	wherein said of Claims 1 To about 800	erystalline aluminosilic erystalline aluminosilic erystalline aluminosilic erystalline aluminosilic to 6 wherein said pro F, a pressure of from	ate zeolite is ZSM-11. ate zeolite is ZSM-12. ate zeolite ZSM-35. ate zeolite is ZSM-38. ate zeolite is ZSM-38. ate zeolite is ZSM-38. beess is conducted at a about 50 psig to about	60	
65	3000 psig, a liquid hourly spac circulation rate of from 0 to al	e velocity of fro	m about 0.1 hr 'to ah	Out 20 hr ' a hydrogen	65	

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oil feedstock is from about 0.1 to about 5. 8. The process of any one of the preceding Claims wherein said gas oil feedstock is a hydrocarbon oil boiling above about 350°F. 9. Process of catalytically dewaxing gas oil substantially as described herein with reference to any one of Examples 3, 5, 6 or 9. 10. Gas oil which has been dewaxed by the process of any one of the preceding Claims.	5
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